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(12)

# EUROPEAN PATENT APPLICATION

(43) Date of publication: 29.08.2001 Bulletin 2001/35 (51) Int Cl.7: C08J 9/12, B32B 5/18 // C08L3/00

- (21) Application number: 01103390.9
- (22) Date of filing: 14.02.2001
- (84) Designated Contracting States: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE TR Designated Extension States: AL LT LV MK RO SI
- (30) Priority: 15.02.2000 IT TO000141
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- (54) Foamed starch sheet
- Materials in the form of foam sheet comprising destructured or complexed starch expanded as a continuous phase, having a density lying between 20 and

150 kg/m3, cell dimensions in the range lying between 25 and 700 µm and with a cell distribution such that 80% of them have a dimension lying between 20 and 400 µm in the absence of stretching.

# Description

## Background of the invention

[0001] The present invention relates to partly-finished products such as sheets of different thicknesses and profile based on destructured and/or complexed starch, expanded by means of an extrusion process, which can be used as such, variously treated, as biodegradable products and which can be formed at the output of the extrusion head or in a subsequent stage, and to products formed there from.

[0002] Starch-based products according to the invention are particularly suitable for use in the packaging sector.

[0003] The use of plastics materials such as polystyrene, polystemane, polystylene and polypropylene has until now dominated in the packaging sector; however, the problems of disposal associated with these products is opening new prospects for starch-based material in that they are blodegradable and from renewable sources, in particular in the foom materials sector.

10004] The state of the art shows various approaches to the formation of foamed starch-based products, However, because of the nature and characteristics of starch 25 it appears at present problematical to succeed in obtaining starch-based foamed products with optimum properties in terms of dimensions and cell distribution, and density of the partly-finished product such as to permit the conversion of the parity-finished product in a regular some answer at an industrial rate into a competitive product as far as weight and performance is concerned, in particular as far as the aspect of ragility of the product at hinge points is concerned. This is particularly true for the preparation of starch-based foams utilised for the stronger of sheets and associated moulded items.

[0005] In particular, no starch-based partly-finished product is yet available on the market with starch in continuous phase, which is able to be shaped using an industrial process, with opimum properties in terms of dismensions, coll distribution and density such as to render the resultant product resilient, in particular in the hinge regions even after successive bending.

[006] In effect, whilst much attention has until now been directed to research and making available various starch-based compositions comprising combinations with various synthetic polymers and additives, the problem of making available extrusion and foaming processes which make it possible to arrive at the production of commed products having well determined properties such as homogeneity of the foamed structure, surface smoothness, and low fragility of the foamed workpieces has received imitted attention.

# Starting composition

[0007] The products according to the invention are obtained from starting compositions supplied to the ex-

truder containing starchy material, water in percentages bying between 4 and 30 percent wt. of the total composition, possibly a thermoplastic polymer and possibly further additives such as plasticizers, lubricants, surfactants, weak acids etc. As far as the components of the starting composition are concerned, the contents of the European Patent Application EP/0 986 811 are incorporated into the present application by reference. [9008] In particular, the present invention relates to foamed, partly-finished products such as sheets of various thickness and profile which can be used themselves as products, and associated shaped on formed products comprising destructured and/or complexed starch as conflictious phase in the partly-finished products comprising destructured and/or complexes.

uct and the finished product itself. [0009] As far as the starch material is concerned this can be both crude and modified starch or a mixture of these. The use of potato, wheat, maize and tapioca starch is preferred. As far as modified starches are concerned, these can be physically and chemically modifled, for example ethoxylated starches, acetate starches, butyrate starches, propionate starches, hydroxypropylated starches, cationic starches, oxidated starches. cross-linked starches, gelatinised starches, starches complexed with molecules and/or polymers able to give "V" type complexes, dextrinated starches and starches grafted with chains such as polyesters, polyurethanes. polyesters-urethanes, polyureas, polyesters-ureas, polyslloxanes, silanes, titanates, fat chains and so on. The preferred chemically or physically modified starches are those with any kind of modification, which have an intrinsic viscosity, measured in DMSO at 30° C, lying between 2 dl/g and 0.6 dl/g, preferably between 1.5 dl/ g and 0.8 dl/g, and more preferably between 1.3 dl/g and 1 dl/a.

[0010] It is intended that flours and meals resulting from the discharge from mill workings lie within the invention.

[0011] The term destructured starch is intended to mean a starch which has been treated thermally above the glass transition temperature and fusion temperature of its components to obtain the consequent disordering of the molecular structure of the starch grains and to render it thermoplastic. Reference is made in this respect to patents EP 118240 and EP 327505.

[0012] Complexed starches on the other hand mean a starch where the amylose component is partially or entirely engaged in the formation of "V" type complexes (single helix structures) which have second derivative X-ray spectral and FTIR characteristics.

[0013] With reference to the thermoplastic polymer, polymers having a melting point or glass transition point lying between 60 and 175 ° C are particularly relevant for the products according to the present invention, and in particular those having such points lying between 70 and 110 ° C.

[0014] In particular usable polymers are selected from:

polymers of natural origin, which can be both modified and non-modified, in particular those derived from cellulose such as cellulose scetate, cellulose propionate, cellulose butyrate and their co-polymers, with a degree of substitution lying between 1 and 2.5; polymers of the alkyl cellulose type, hydroxyalkyl cellulose, carboxyalkyl cellulose, in particular carboxymethyl cellulose, nitrocellulose and chibosane, pullulan or caselin and caselinate, zein, soya protein, alginic acid and alginates, natural rubbers. polyasporates, glutien;

biodegradable polymers of synthetic or fermentative origin, in particular polyesters, such as polymers or co-polymers, of C2-C24 aliphatic hydroxyacids, or their corresponding lactones or lactides, in particular polymers of lactic acid having various D/L lactic acid ratios, and preferably with a D-lactic content comprised between 4 - 25% mole, co-polymers of polylactic acid with aliphatic polyesters and aromatic-aliphatic polyesters, polycaprolactone, polyvalerolactone, their co-polymers and polyesters derived from difunctional acids and aliphatic diols, aliphatic-aromatic polyesters, in particular copolymers of the alkylene-terephthalate adipate type whether treated or not with chain extenders, prefer- 25 ably with quantities of terephthalic acid less than 40 mole percent, preferably less than 30% mole, epoxy resins in general and bisphenolic resins in particu-

polymers able to interact the starch to form complexes, that is to say polymers which contain hydrophilic groups intercalated with hydrophobic sequences, for example, sinylenewiny alcotate co-polymers, earlyic esters, ethylene ecrylic ester co-polymers, conjuines, ethylene exitylic ester co-polymers, ethylene exit, co-polymers, ethylene ethic, co-polymers having alcoholic and carboxytic functional groups aliphatic polyseters end/or aliphatic-aromatic polyseters, epoxy resins including those containing bisphenol resins;

polymers forming hydrogen bonds with starch, in particular, polyvinyl alcohols of varying degrees of hydrolysis, possibly modified as acrylates or methacrylates and polyvinyl alcohols preliminarily plasticised or modified for the purpose of lowering the melting point.

[0015] Preferred thermoplastic polymers are the polyinyl alcohols, co-polymers of an olefinic monomer, preferably ethylene, with a monomer chosen from vinyl alcohol, vinyl acetate, acrylic acid and methacrylic acid, alphatic polysers derived from caprolactone, polyalk-ylenesuccinates, polymers of azelaic acid, sebacic acid, senssilic acid and their co-polymers, aliphatic polyamides, polyalkylenesebacates, polyalkyleneazelates, polyalkyleneazessilates, in particular with diols comprised between  $C_{\mathcal{P}}C_{12}$ , polyesters containing dimeric

acids, aromatic-aliphatic polymers of the polyalkylene terephthalate adipate type and the epoxy resins, particularly with bisphenolic groups.

[0016] The extruded feamed partly-finished product according to the invention further preferably contains a nucleating agent. The use of a suitable nucleating agent in fact makes it possible to increase the homogeneity of the cells of the sheet. The quantity of nucleating agent utilised in the course of the process depends on the process conditions and the desired morphology for the extruded, partly-finished product. Proferably, the quantity of nucleating agent with respect to the starting composition lies in the range between 0.5 and 7 % and more preferably between 1 and 5 %.

[0017] Usable nucleating agents are, for example, inorganic compounds such as talc (magnesium silicate). calclum carbonate, sulphates such as sodium and barium, titanium dioxide etc, possibly surface treated with adhesion promotors such as silanes, titanates, etc. Organic fillers and fibres such as wood powder, cellulose powder, grape residue, bran, maize husks, other natural fibres in concentrations between 0.5 and 20% may also be utilised. Further, substances able to be dispersed and/or to be reduced in lamellas with submicronic dimensions, preferably less than 500 µm, more preferably less than 300 µm, and even more preferably less than 50 µm may be utilised in order to improve stiffness, water and gas permeability, dimensional stability, Particularly preferred are zeolites and silicates of various kind such as wollastonites, montmorillonites, hydrotalcytes functionalised with molecules able to interact with starch. Particularly preferred are submicronic particles of complexed starch also with specific functional groups Introduced by virtue of sllanes, titanates and other.

[0018] The starting compositions can moreover contain suitable additives such as lubricating agents and/or dispersants, flame retardants, colorants, plasticising agents, fillers etc. In particular, food oils such as paim. maize, soya, sunflower oil are particularly good, as are fatty acids from C12 to C22 and their glycerides with various degrees of substitution and in particular synthetic hydrogenated fats or fats of animal origin which are solid at least at ambient temperatures and, preferably, above ambient temperature to improve the moisture resistance and reduce wetability. It is also possible to use weak acids such as lactic, tartaric, citric acid etc to regulate the viscosity of the starch during the extrusion, and plasticizers such as glycerine, sorbitol, mannitol, pentaerithritol, and derivatives thereof, esters of citric acid and their derivatives

[0019] The starting composition can be supplied directly to the extruder or can be supplied in the form of preliminarily extruded or pelletised granules.

# Process for the production of the sheet

[0020] The foamed, partly-finished product according

to the Invention is prepared by means of a process of extrusion of the basic starch composition effected by means of particular extruders such as slow twin screw extruders or two single screw tandem extruders in cascade or their combination, in such a way as to guarantee significantly long dwell times for the purpose of optimising the viscosity of the starchy material and the homogenisation of the nucleating agents and mixing of the foamting agents in the molten mass. In particular the use of a slow twin screw extruder is preferred.

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[0021] In the performance of the extrusion process the extrusion temperature can vary as a function of the particular formulation and the desired properties of the partiy-finished product and the finished product. The temperature control of the molent mass is therefore significant for obtaining products with specific characteristics.

[0022] The temperature of the molten mass in the course of the extrusion process can generally vary along the profile of the screw from 50 to 230° C, preferably between 60 and 210° C and more preferably between 70 and 200° C.

[0023] The foarning of the thermoplastic products ecording to the invention is achieved by the use of a suitable mixture of physical foarning agents which can also contain chemical foarning agents. In particular the use of  $Co_2$  in gaseous form is preferred, in combination with water or  $CO_2$  in gaseous form in combination with water or the physical and chemical foarning agents. Among the chemical foarning agents can be taken into consideration, among others, citric acid, bicarbonate and their combinations.

[0024] The foaming agent is preferably supplied to a region of the extruder in which the starting composition supplied to the extruder is present in the molten state. In particular the foaming agent is supplied to an advanced region of the extruder in such a way that the extrusion process is not altered by the phenomenon of regurgitation of the molten mass towards the extruderfeed zone.

[0025] The CO<sub>2</sub> is supplied in concentrations greater than 0.4%, preferably greater than 0.8%, with respect to the total composition fed to the hopper, to a region where the melt is at a temperature lying between 100° C and 200° C, preferably between 130° C and 100° C. 45 The mixture of CO<sub>2</sub> and H<sub>2</sub>O and the specific concentrations are determining factors for the low density and the cell structure.

[D026] The quantity of CO<sub>2</sub> can vary in a range lying between 0,4% and 10% by weight, preferably between 50,8% and 7% and more preferably between 1% and 4% by weight. The CO<sub>2</sub> is added to the melt. The total water content of the composition fed to the hopper of the extruder for the expansion is lying between 4% and 30% by weight, preferably between 1% and 12% by weight, preferably between 10% and 18% by weight.

[0027] The extruder can be completed by extruder heads of the flat or tubular type; tubular heads are par-

ticularly preferred.

10028] Preferably the configuration of the head of the extruder is such as to guarantee a homogeneous supply to the nozzle. With the starting compositions of the foam sheet according to the invention this problem is relevant since small variations of sheer rate can generate significant variations in the local viscosity, with consequent alterations in the foaming process and therefore manifest irregularities in the sheet in terms of thickness thereof, cell dimensions, presence of preferratial flow etc.

[0023] The head of the extruder is therefore proferably configured in such a way as to cancel the elastic memory of the material and, at the same time, not create any foam before the entry of the material into the nozzle. The preferred extrusion shear rate ranges for the sheet are between 500 and 50,000 sec<sup>-1</sup>, preferably between 800 and 40,000 sec<sup>-1</sup>, and more preferably between 900 and 55,000 sec<sup>-1</sup>.

[0030] At the output from the extrusion head, and before the forming process, the foamed sheet according to the invention can be laminated with layers of non-woven fabric, textile, paper, biodegradable and non-biodegradable films, or aluminium. As far as the non-woven or textile fabrics are concerned these can be made of natural fibres, such as, for example, fibres of jute, cotton. wool, fibres based on polysaccharides such as, for example, cellulose acetate, starch acetate, viscose etc. or fibres produced from biodegradable polymers and in particular aliphatic polyesters such as polylactic acid. polycaprolactone, polyalkylene carboxylate with dialcohols and diacids selected from the linear range Co-Cta and/or cycloallphatic, aliphatic-aromatic polyesters, in particular of the family of terephthalate polyalkylene adipates and their co-polymers, particularly with a terephthalic acid content less than 55% with respect to the amount of terephtalic acid + adipic acid, polyamides in particular based on caprolactam, aliphatic amines etc, aliphatic polyurethanes, polyester-urethanes, polyurea. and epoxy resins. The above biodegradable polymers can be utilised also in the form of films for lamination or coating.

[0031] The films are generally coupled to the sheet through temperature and/or the application of suitable biodegradable achesives based on polymers, lactic acid, polymerthenes, polyrinyl accetates and polyvinyl alcohol, proteins such as caselin and glutens, starches and other polyseccherides, hot melts particularly based on aliohaltic polysecs.

[0032] The films can be obtained by casting or bubble film-forming and can be co-extruded with an adhesive surface for the foam support. Films with a melting point greater than 60° C, preferably greater than 80° C, and more preferably greater than 100° C, are preferred.

[0033] The partly-finished product coupled to film can be used unformed, as the sheet as such, or in a foaming or shaping process to form finished products.

[0034] For the coating it is possible to utilise emulsions, solutions or dispersions of the type described in European patent EP 896612 for the treatment of expanded particles, considered included within the present invention. Natural and synthetic waxes can also be utilised, with melting points up to 120° C depending on the application. In this case the treatment can be before or after the forming or shaping stage.

[0035] The toam sheet may also be co-extruded with expanded layers of other starch based materials so to have differentiated properties between the inside and the outside of a multiplayer or it may be co-extruded with layers of materials having lower hydrophilicity such as the polyesters above mentioned for the laminated films. [0036] The foam sheet according to the invention must be obtained starting from a homogeneous molten mass in which the nucleating agenits and the gas and or vapours are homogeneously dispersed throughout the molten mass. For this the dwell times in the extruder must lib between 5 and 40 minutes, prefarably between 10 and 35 minutes, and more preferably between 16 and 25 minutes.

[0037] The foam sheet can be controlled in thickness by the extrusion conditions and calendering.

[0038] In the case of tubular sheet the head can be provided with air or steam blowing systems from within, as in the case of bubble film-forming, or or enlation by 2s air or steam blowing to distend the sheet and give it a biaxial stretch, avoiding or regulating the formation of waves. The sheet can have a thickness lying between 0.5 mm and 15 mm, preferably between 1,0 mm and 10 mm. The thickness of the parity-finished product can be achieved by stretching and celendering the sheet.

[0039] The foam sheet may be corrugated and the corrugations may be exploited in order to increase the cushion properties of the foam. The corrugations may have different width and height. The height, which corresponds to the thickness of the resulting panel, may be about the double of the foam sheet thickness. The frequency of the corrugations may reach 350 per linear me-

[0040] Sheets of this type can be combined together in multi layers, forming different geometries for different products in the packaging sector such as sheets and expanded blocks of high resilience, corners or protection containers. Specific examples of the sectors suitable for application are those of electrical domestic equipment, electronic products, the food sector, pharmaceuticals, design and furniture, mail order, and envelopes for couriers. The sheets can be utilised also in combination with other supports to form multi layers mixed with wood, paper, cardboard, textiles of natural and syn- 50 thetic fibres, aluminium and other metals. In particular, the products of this type can be directly anchored to the piece to be packaged, exploiting their characteristic adhesiveness upon moistening or, preferably, with hot melts or melts to be sprayed.

[0041] Products obtained from coupling sheets or formed products can be protected by an external film to increase performance.

[0042] Products can also be rolls and tubes obtained by winding and gluing sheets, or by directly extruding tubes. Rolls and tubes can be utilised as supports for tollet paper, kitchen paper or other types or may be used as protection for cylindrical things such as bottles and others.

## Process for forming the sheet

 [0043] Forming can be achieved by a continuous process or by a batch process.

[0044] The production of expanded products according to the invention by means of continuous processes provides for the extrusion/calendering phase, a possible

conditioning phase and the forming phase to be consective. The production of foamed products according to the invention by means of batch processes provides for the axtrusion/calendering phase with winding of the sheet into color or collection in sheets; the partly-finished products can them be conditioned and formed in a second phase.

(0045) In a continuous fearing process it is envisaged that the partly-finished product in the form of sheet from the extruder would be maintained at a temperature not less than 40° C, and preferably not less than 80° C and having a water content lying between 6 and 30% by weight, preferably between 10 and 25% by weight and more preferably between 15 and 20% by weight. The temperature of the sheet must not exceed 150° C and preferably 100° C.

[0046] In particular, if synthetic components are present, the forming temperature must be close to the glass transition temperature or the melting point of the thermoplastic polymer.

50427 It is also possible to form the partly-finished product by a batch process by subjecting it to a preliminary conditioning process for the water content and temperature range referred to above for continuous processes.

[0048] The conditioning stage can immediately precede or be coincident with the forming station.

[0049] Products even of complex form, provided with particular, even aesthetic, characteristics such as, for example, the claim shell illustrated in Figure 1, can be 5 obtained with a forming process at ambient temperature, and in any event at temperatures not greater than 100° C, between abutting male and female mould to define the maximum level of compression and the final minimum thistness of the product.

[0050] A die for the forming of the clarm shell of Figure 1 with the foam sheat according to the present invention is illustrated, as a way of example, in Fig. 2. Male (10) and female (11) may be designed in such, a way they do not got in touch just next to the lateral walls of the clarm shell. Such room between male and female allows the sitepring of the foam sheet without tears during

[0051] The process forming the subject of the inven-

the forming

tion, together with the characteristics of the partly-finished product generally allow forming cycles less than 20 seconds, preferably less than 10 seconds and more preferably less than 7 seconds. With reference to the forming process, the parameters relating to the water content and temporature are critical for the achievement of a good formability of the partly-finished product. The loss of water vapour from the partly-finished product at the outlet from the extruder nozzle in fact makes it necseasing to exercise a strict control on the level of removal of water for the purpose of avoiding both phenomena of collapse and phenomena of excessive driving.

[0052] Forming can take place in moulds or dies at ambient temperature on expanded but unopened tubular sheets. This system makes it possible simultaneously to mould two layers of sheet per mould, limiting the problems of drying of the sheets. If the tubular sheets are conveniently offset from one another it is possible to obtain a surface of the product having an aspect similar to the surface weeve of a fabric.

[0053] Forming is normally conducted on an opened tube. In this case the water content is regulated by utilising a conditioning station which uses steam.

# Characteristics of the sheet

[0054] The material which constitutes the partly-finlished product or fearned sheet forming the subject of the present invention has an intrinsic viscosity in DMSO at 30° C lying between 1.5 and 0.3 d/g, preferably lying between 1.2 and 0.4 d/g and more preferably between 1 and 0.8 d/g.

[0055] In expanded partly-finished products according to the invention the cell dimension can vary in a range lying between 25 and 700  $\mu m$  and preferably between 40 and 600  $\mu m$  (as determined by microscopic inspection).

[0056] The expanded partly-finished product has closed cell morphology in which the cells are substantially non communicating with one another, which is different from the open cell morphology in which the cells are larquely interconnected with one another.

[0057] The partly-finished product can have a density lying between 20 and 150 kg/m³, preferably lying between 25 and 100 kg/m³, more preferably between 30 45 and 70 kg/m³.

[0058] The foam structure of the sheet is characterised by a cell distribution in which 80% of the cells present, in the absence of stretching, have a dimension lying between 20 and 400 µm, preferably between 25 and 300 µm and more preferably between 30 and 200 µm.

[0059] When a stretch is applied to the sheet the cells can, however, be subjected to an orientation with thinning of the wall.

[0060] Also within the scope of the present invention is a sheet with optimised resilience properties, a density characteristic lying between 30 and 70 kg/m³ and with

an average cell dimension between 80 and 120µm. 10061] Products forming the subject of the present invention are principally used in the food packaging sector and in particular as trays for food with a lifetime of the order of 30 days, for the packaging of meat, milk products, vegetables, eggs and fruit, holders for packages of glass, plastics or metal of very small dimensions, containers for fast food such as containers for hamburgers, potato chips and similar products; multi compartment containers for foods, known also as lunch boxes, cups for coffee and other hot or cold drinks for fast food and meals.

[0062] The formed products of the present invention are also used as containers for objects of small weight such as multicompartment trays for portable telephones and small electrical domestic appliances in particular, with mechanical properties such ast ox oxid phenomena of abrasion encountered with containers of pressed paper etc.

20 [0063] In the case of food applications where liquids at high or low temperatures are to be expected, the containers can be co-extruded or coupled to another layer of foam or polyester film and/or cellulose acetate and/ or starch or other polymer resistant to liquids at the tem-25 perature which will be experienced in use. In particular films of aromatic-aliphatic polyester type can be utilised and, specifically, polyalkylene terephthalate adipates. alkylene butyrates, polyalkylene succinates, polyalkylene sebacates, polyalkylene azelates, polycyclic alkylene dicarboxylates, in particular polyhexyldimethyldicarboxylates, olycyclohexyldicarboxylates. If it is necessary to absorb liquids as in the case of packaging for meat it is possible to consider the use of superabsorbent material which can inserted directly into the sheet, applied to the surface or in intermediate lavers between two shells welded together or under the film which makes the tray impermeable.

[0084] Also to be considered the subject of the present invention are products formed for ovens and microwaves, possibly characterised by treatments with water-repellent coatings to avoid drying of the container during the cooking phase.

#### Characteristics of the formed products

[0065] Formed products according to the invention have a closed cell structure with a relatively low density lying between 40 and 400 kg/m³, preferably between 45 and 200 kg/m³ and more preferably between 50 and 150 kg/m³.

[0066] Products formed according to the invention further have good properties of flexibility, in particular in the hinge region, thanks to the fine and homogeneous morphology of the cells. Such products also have a very good uniform surface.

[0067] Hinges, such as for examples the one numbered as 12 in Figure 2, can be produced in products obtained in the forming phase, by forming ribs of the type used for cardboard hinges, are resistant to at least ten (preferably > 20) consecutive opening at 180°/closing cycles at 35%. RH and 23°C without breakage, using about 2-4 seconds for each opening at 180°/closing operation, and preferably at least 100 consecutive opening and closing cycles at 40%. RH and 23°C without breaking, using about 2-4 seconds for each opening/closing cycles.

[0068] The good properties of flexibility can be tested also with a dynamometer with a climinatic cell adapted to 16 adjust the temperature and relative humidity at the above values. Samples of 25 x 10 cm with an hinge at the middle of their length can be submitted to opening/closing cycles from 0 to 160° with a velocity of in the range of 3000 - 10,000 mm/min of the mobile bar of the 40 ynamometer.

## Examples

[0069] The Invention is further illustrated by means of the following examples provided by way of illustrative and non-limitative example of the invention itself.

# Example 1

[0070] A mixture was prepared having the following composition:

- 88.9% of destructured potato starch with an intrinsic viscosity in DMSO at 30°C of 1.1 dl/g and the water 30 content of 14%.
- 8.9% by weight of polyvinylalcohol
- 1.8% by weight of talc
- 0.35% by weight of glycerol
- 0.36% by weight of loxial G10
- 2% by weight of water.

[0071] The composition was supplied to a slow twin screw extruder with co-rolating screw having a diameter (a) = 113.8 mm and L/D ratio = 19:1. At the end of the extruder was mounted an extrusion head for a tubular sheet with a diameter of 100 mm and lip opening of 0.5 mm. The dwell time of the mel in the extruder was about 20 minutes.

[0072] In addition to the water contained in the feed mixture, a further 1% by weight of  $CO_2$  was also added to the molten mass as a further expansion agent, at a feed pressure equal to 37 bar. The  $CO_2$  was introduced at the level of the eleventh diameter of the screw.

[0073] The operating conditions were as follows:

- BPM: 16
- Temperature profile (°C): 95/120/120/150/ 180/185/190/197
- Feed rate: 54 kg/h
  - Lip shear rate: 912 sec -1

[0074] The foamed sheet obtained had a density of

56 kg/m³ and a cell dimension lying between 40 and 170 μm, the average value of the cell dimension was 81 μm. [0075] The Intrinsic viscosity of the material constituting the sheet, taken in DMSO at 30°C, is 0.68dt/g.

The sheet was wound in a coil.

## Example 2

[0076] A mixture was prepared having the following composition:

- wheat starch 34.4% (12% H<sub>2</sub>O)
- potato starch 34.4% (16% H<sub>2</sub>O)
- polyvinylalcohol 13.5%
   H<sub>2</sub>O 17.4%
- Monoglyceride oleic acid 0.3%

[0077] This mixture was supplied to a twin screw APV 2080 extruder having a diameter (d) = 80mm and L/D ratio = 40. It was operated in the following conditions:

- BPM: 28
- Temperature profile: 50/75/75/180/180/170/ 170/175/175/185/185/155/155/154/120 Degassing was regulated in such a way as to maintain in the granules a total water content of about 14.5%. The intrinsic viscosity of the pellets was 1.98 dl/g.

9078) The granules thus obtained were mixed with 3 2.5% of talc having an average particle cliemeter of about 1.5 µm and supplied to a slow tivin screw extruder with co-rotating screws having a cliemeter (d) = 113.8 nm and L/D ratice = 181 · with an extrusion head for tubular sheet of 100 nm in diameter and 0.4 mm of lip separation, operating in the following conditions:

- RPM; 14
- Temperature profile (°C): 90/120/120/140/ 165/165/170/186/186
- Feed rate : 50 kg/h
- Shear rate: 1360 sec-1

[0079] To the molten mass was added, as a further expanding agent, CO<sub>2</sub> in quantities equal to 1.5% by weight at a feed pressure equal to 40 bar.

The tubular sheet obtained had a thickness equal to about 3 mm a density of 70 kg per m³ and an average cell dimension equal to 90 µm (minimum/maximum cell dimension = 102:90 µm). The water content of the sheet was equal to about 1.8% by weight and the intrinsic viscosity of the material constituting the sheet was  $m=1.1\,$  d/g.

# Example 3

[0080] The tubular sheet obtained by the example 2 was opened by subjecting it to a calendering and steam conditioning process until it had a water content in the

sheet equal to 15 %. The forming was achieved by means of a suitable mould, such that illustrated in figure 2, of the male-female type for hinged trays of the clarn shell type suitable for fast food products.

[0081] Forming was conducted with dies at ambient temperature on the sheet maintained at a temperature about 80°C by applying a pressure of 6 kg/cm<sup>2</sup>. The moulding cycle was about 6 seconds and the product thus obtained had a thickness equal to about 1.6 mm and a density in the bottom wall of 165 kg/m3.

[0082] In particular, the product obtained was constituted by two asymmetrical valves having a length of 12.5 cm connected by a hinge 10 cm wide. This hinge zone had particular properties of mechanical strength. After 20 successive bendings for a time of 3 seconds (corresponding to about 5000 mm/min) for opening/closure cycles at 35% RH and 23° C it continued to perform its function.

[0083] The product obtained also had a very smooth surface constituted by super-imposed flattened ribs which confer on the product a pleasing aesthetic aspect.

#### Example 4

[0084] The colled sheet obtained according to example 1 was maintained at a water content of 14%. To the sheet was applied a film of 14 µm of polybutyleneterephthalate-adipate containing 33% by mole of terephthalate with an intrinsic viscosity in THF of 1.1 dVg. The sheet with the applied film was brought to 80°C and formed in the mould described in example 3. The container obtained was resistant to water at 80°C for an hour, the time necessary for the temperature to fail from 80°C to 20°C without any collapse or soaking of the foamed starch container.

## Example 5

[0085] As for example 4, with the single difference of having applied a film of polyethylene sebacate. The tray was perfectly resistant to water without becoming saturated and/or collapse of the starchy product.

#### Example 6

[0086] As for example 4, with the exception of the application of a non-woven fabric of viscose of 30 g/m2, in place of the polyester film.

#### Example 7

[0087] As for example 4, with the exception that the polyester film was replaced by a foamed sheet of polyethylene sebacate of a density of 80 kg/m3 and a thickness of 300 µm.

## Example 8

[0088] As for example 4, with the exception that the film was applied to both sides. The resultant tray was utilised for packaging trials of beef. The results related to the mechanical properties and to the preservation of the meat were comparable to the ones observed for trays made with expanded polystyrene.

## Example 9

[0089] The tubular sheet obtained according to process of example 1 was formed directly in the conditions of example 3 with a male/female mould in the form of a tray 2.5 cm deep and 15 x 12 cm, to form a double container with a thickness of about 3 mm. The double container was positioned between two films of the type described in example 4, of 10 µm which were welded together forming a bag within which the tray was contained. The film was heat shrunk to form a compact and impermeable product for meat.

# Example 10

[0090] A mixture was prepared having the following composition:

- 74.3% by weight of potato starch (H<sub>2</sub>O 16%)
- 10.0% by weight of Ecoflex EBX 7000 (BASF) 0.3% by weight of Loxiol G 10 F
- 15.4 by weight of water.

The composition was supplied to a twin screw extruder APV 2030 with (d) = 30.0 mm and L/D = 40. The operating conditions were as follows:

- RPM:
- Temperature profile (°C): 30/100/100/150/ 160/150/140/130/110 x 8

[0091] The degassing step was adjusted so as to have in the pellets a water content of about 13.5-14.5%. [0092] The pellets were then mixed with 2.5% of talc, with particles having mean diameter of 1.5 um, and subsequently fed to a slow twin screw extruder with co-rotating screws having a diameter (d) = 113.8 mm and L/ D ratio = 19:1. At the end of the extruder was mounted an extrusion head for a tubular sheet with a diameter of 100 mm and lip opening of 0.1 mm. The operating con-50 ditions were as follows:

RPM: Temperature profile (°C): 90/120/140/180/

210/210/210/195/196 Feed rate :

> shear rate : 31531 sec -1

fed composition, of  $\mathrm{CO}_2$  was also added to the molten mass as a further expansion agent, at a feed pressure equal to 40 bar. The obtained foamed sheet had a thickness of about 5 mm, a density of 81 kg/m³ (calendered) and a everage value of the cell dimension of 85  $\mu$ m (cell dimension by fing between 35 and 188  $\mu$ m).

#### Claims

- A partly-finished product, in particular in the form of a foam sheet material, comprising destructured or complexed starch foamed as a continuous phase, having a density lying between 20 and 150 kg/m³, cell dimensions in a range lying between 25 and 700 μm with a cell distribution such that 80% of them have, in the absence of stratching, a dimension lying between 20 and 400 μm.
- 2. A parity-finished product, in particular in the form of 20 a foam sheet material, according to claim 1 having a density lying between 25 and 100 kgm³ and ceil dimensions in a range lying between 40 and 800 µm and with a ceil distribution such that 80% of them have, in the absence of stretching, a dimension lying between 25 and 300 µm.
- A partly-finished product, in particular in the form of a foam sheet material, according to claim 2 having a density lying between 30 and 70 kg/m³ and with a cell distribution such that 80% of them have, in the absence of stretching, a dimension lying between 30 and 200 µm.
- A partly-finished product, in particular in the form of foam sheet, according to claim 3 having a density lying between 30 and 70 kg/m<sup>2</sup> and average cell dimensions lying between 80 and 120 µm.
- A partly-finished product according to any of claims 40
  1 to 4, in which the starch is natural or modified starch or a mixture of these.
- A parity-finished product according to claim 5 in which the natural or modified starch is derived from 45 potato, wheat, maize and taploca.
- 7. A parily-finished product according to claim 5 wherein the modified starch is physically or chempalary modified, particularly ethoxylated starches, pacotate starches, butyrate starches, propionate starches, hydroxypropylated starches, cationic starches, oxidated starches, cross-linked starches, gelatinised starches, starches complexed with not-ceules and/or polymers able to give "V" type complexes, dextrinated starches and starches grafted with chains such as polyesters, polyurethanes, polyesters-ureath, polyesters

polysi-loxanes, silanes, titanates, fat chains.

- 8. A parity-finished product according to any of claims 1 to 7, able to form products with hinges obtained in a forming phase capable of resisting at least ten consecutive opening/closing cycles at 35% RH and 23°C without breaking, by using 2 - 4 seconds for each opening and closing operation.
- 9. A partly-finished product according to any of claims 1 to 8, in which the material from which the foam sheet is made has an intrinsic viscosity in DMSO at 30°C lying between 1.5 and 0.3 dl/g.
  - A partly-finished product according claims 9 in which the intrinsic viscosity in DMSO at 30°C is lying between 1.2 and 0.4 dl/g.
  - A partly-finished product according to claim 10, in which the intrinsic viscosity in DMSO at 30°C is lying between 1.1 and 0.6 dl/g.
  - A partly-finished product according to any of claims from 1 to 11, containing one or more thermoplastic polymers with a melting point lying between 60 and 175°C.
  - 13. A parthy-finished product according to claim 12 in which the thermoplastic polymer is a polymer of natural origin which can be modified or non modified, in particular derived from cellulose as cellulose acetate, cellulose proplonate, cellulose buyrate and their co-polymers, with a degree of substitution lying between 1 and 2.5; polymers of the alkyl cellulose, hydroxy alkyl cellulose, carboxy alkyl cellulose, hitrocellulose and chitosan pullulan or casein and casinate, zein, soya protein, alginic acid and alginates, natural rubbers, polyaspartates; glutnes, dextrens.
  - 14. A partly-finished product according to claim 12 in which the thermoplastic polymer is a biodegradable polymers of synthetic or fermentative origin, in particular polyesters of the type including polymers or co-polymers of C2-C24 aliphatic hydroxy acids, or their corresponding lactones or lactides, in particular polymers of lactic acid having various D/L lactic acid ratios, co-polymers of polylactic acid with aliphatic and aliphatic-aromatic polyesters, polycaprolactone, polyvalerolactone, their co-polymers and also polyesters derived from difunctional acids and aliphatic diols, aliphatic-aromatic polyesters, in particular co-polymers of the type including alkaline-terephthalate adipate treated or not with chain extenders, preferably with quantities of tereftalic acid less than forty mole percent, epoxy resin in general and bisphenolic resin in particular.

- 15. A parily-finished product according to claim 12 in which the thermoplastic polymer is a polymer containing hydrophilic groups intercalated in hydrophobic sequences such as, for example, ethylene-viny-lalcohol co-polymers, orbitheen vinylacetae co-polymers, acrylic esters, acrylic ethylene-ester co-polymers, co-polymers of ethylene with unsaturated acids such as acrylic acid, methacrylic acid, croton-ic acid, itaconic acid, co-polymers with hydrophilic untia with a functional alcoholic a carboxylic group in aliphatic polyesters and/or aromatic-aliphatic polyesters, epoxy resins including resins containing bisphanols.
- 16. A parity-finished product according to claim 12 in 15 which the thermoplastic polymer is a polymer able to form hydrogen bonds with the starch, in particular polyvinyl alcohol with various degrees of hydroylas, possibly modified with explicates or methacrylates, polyvinyl alcohol preliminarily plastisized or modified for the purpose of lowering its mething point.
- 17. A parity-finished product according to Claim 12 containing polymers such as polyvinylalcohol, copolymers of an olefin polymer, preferably ethylene, with a monomer chosen from vinyl alcohol, vinyl acetate, acrylic acid and methacrylic acid, aliphatic polyesters such as caprolactone, the polyalkylene succinates, the polymers of azellac acid, sebacic acid brassilic acid and their co-polymers, aliphatic polyamides, polyalkylenesebacates, polyalkylene-azelates, polyalkylenebrassilates, in particular with diols comprised between 0<sub>2</sub>-C<sub>13</sub>-polyesters containing dimeric acids, aromatic-aliphatic polymers of the polyalkylene terspithalae acipate type and the aspony resins, particularly with bisphanolic groups.
- 18. Partly-finished products according to any of Cleims from 1 to 17, containing nucleating agents for the starting composition in concentrations lying in the range from 0.05 to 10% by weight, preferably between 0.5 and 7% and more preferably between 1 and 6%.
- A parily-finished product according to Claim 18, in which the nucleating agent is constituted by inorganic compositions such as talc (magnesium silicate), calcium carbonate, sulphates of sodium and barium, fitanium dioxide, possibly surface treated with adhesion promoters such as silinare, stitanates.
- A partly-finished product according to any of Claims from 1 to 19 containing organic fillers and fibres such as wood powder, cellulose, grape residue powder, bran, malze husks or other natural fibres in concentrations between 0.5 and 20%.
- 21. A partly-finished product according to any of Claims

- from 1 to 20, containing nucleating agents, lubricants and/or dispersants and plasticisers.
- 22. A parthy-finished product according to any of Claims from 1 to 21 containing alimentary oils such as paim oil, maize oil, soya oil, sunflower oil, C<sub>12</sub> to C<sub>22</sub> fathy acids, their glycerides with various degrees of substitution, and in particular hydrogenated fats of animal or synthetic origin which are solid at least at ambient temperatures, no improve the moleture resistance and reduce the wetstallily by water.
- 23. A parity-finished product according to any of Claims from 1 to 22 containing weak acids such as lactic acid, tartaric acid, citric acid to regulate the viscosity of the starch during the extrusion process.
- 24. Products and partly-finished products obtained from the partly-finished products of Clalms 1 to 23, obtained by lamination with layers of non-woven fabric, woven fabric, paper, biodegradable and nonbiodegradable films or aluminium.
- 25. Products and partly-finished products according to claim 24 produced by lamination with non-woven fabric or woven fabric of natural fibres, such as for example fibres of jute, cotton, wool, fibre based on polysaccharides such as, for example, cellulose acetate, starch acetate, viscose etc. or fibres produced starting from biodegradable polymers and in particular aliphatic polyesters such as polylactic acld, polycaprolactone, polyalkaline carboxylates with dle alcohols and die acids selected from the linear range Co-Cto and/or cycloalophatic, allphatic-aromatic polyesters, in particular from the family of adipated terraphelate polyalkalines and their co-polymers, polyamides, in particular based on caprolactane, aliphatic amine etc. aliphatic polyurethanes. polyester-urethanes, polyurea, and epoxy resins.
- 26. Products and parily-finished products according to claim 24 coupled with films constituted by biode-gradable polymers and in particular aliphatic polyesters such as polylactic acid, polycaprolactone and/or cycloalyphatics, polyalkaline carboxylates with dialcohols and diacids selected from the linear range Cg. Cg. aliphatic-aromatic polyesters, in particular from the family of adipated terephthalate polyalkylenes and their co-polymers, polyamides, in particular based on caprolactam, aliphatic amines etc, aliphatic polyurethanes, polyester-ure-thanes, polyuress, poby reisas, bobalned by blown extrusion, co-extrusion and/or casting.
- 27. Products and partly-finished products obtained from the partly-finished products of Claims from 1 to 23, by way of coating with emulsions, disper-

sions, solutions, hot melts of biodegradable polymers and in particular aliphatic polysters such as polylactic acid, polycaprolactone, polyalitylene carboxylates with dialcohols and diacids selected from the linear and/or cyclealophatics range C<sub>2</sub>·C<sub>13</sub>, allphatic-aromatic polyesters, in particular from the family of adipated polyalitylene terephthalates and helir co-polymers, polyamides, in particular based on caprolactane, aliphatic amines etc, aliphatic polyurehanes, polyester-urethanes, polyureas, epoxy resine.

- 28. Products and partly-finished products according to Claim 26, in which the films are coupled to the partly-finished products by temperature and/or the application of suitable biodegradable adhesives based on polymers of lactic acid, polyurethanes, polyvinylactates and polyvinylalcohols, proteins such as casein and gluten, starches, dextrins and other polyseacharides.
- 29. Products and parity-finished products according to Claims 26 and 28, in which the films can be obtained from cast and bubble film-forming and can be coextruded with an adhesive surface for the foamed 25 support.
- 30. Products and parity-finished products according to Claim 29, in which the films have a meiting point greater than 80° C, preferably greater than 80°C and more preferably greater than 100°C.
- A sheet according to Claim 26 or Claim 28 form able as a non-laminated sheet.
- 32. Products and partly-finished products obtained from the materials of Claims 1 to 23, treated with natural and synthetic waxes with melting points up to 120°C in depending on their various applications.
- 33. A process for the production of foam sheet by extruder comprising the steps of:
  - supplying to an extruder starch with an intrinsic viscosity lying between 2 and 0.6 dl/g in the presence of water in proportions from 8 to 30% by weight of the total composition, in quantities such as to permit the starchy component constitute the continuous phase of the material, possibly a natural or synthetic thermoplastic polymer and further additives such as plasticleers, lubricants, nucleating agents, surfactants, weak acids and fillers.
  - complete melting of the starchy mass
  - introduction of CO<sub>2</sub> in quantities lying between 0.4 and 10%, preferably between 0.8 and 7% and more preferably between 1.0 and 4% by weight into the melt at a temperature lying be-

- tween 100 and 180°C, preferably between 120 and 160°C; and
- working the melt for between 5 and 40 minutes to homogenise the distribution of the mixture of expanding agents, water and CO<sub>2</sub> and, possibly, chemical expanding agents such as citric acid and bicarbonate, and to adjust the viscosity of the composition to between 1.5 and 0.3 dVn
- 34. A process according to Claim 33 in which the extrusion of the melt takes place through a filtar of tubular head able to impart to the melt shear rates comprised between 500 and 50,000 sec<sup>-1</sup> preferably between 800 and 40,000 sec<sup>-1</sup> and more preferably between 900 and 35,000 sec<sup>-1</sup>.
- 35. A process according to Claim 34 in which the extruded tribular sheet is blown with air or steam to impart by blazial stretch, confer smooth surfaces and hold the sheet at the desired moisture point, opened, calendered, possibly further conditioned and wound on a coil.
- 36. A process according to Claim 34 in which the tubular sheet is blown with air or steam to distend the sheet itself and hold it at the desired moisture point, opened calendered and cut into flat sheets.
- A process according Claim 34, in which there is produced a partly-finished product in the form of a tube which is calibrated, conditioned and then collected.
- 38. A process for forming partly-finished products according to any of Claims from 1 to 32 which comprises:
  - conditioning the product or partly-finished product to a water content between 6% and 30% preferably between 10% and 25% and more preferably between 15% and 20% and at a temperature between 40 and 120 °C and preferably between 40 and 100 °C.
  - forming in a male-female impact mould between ambient temperature and 80° C
    - possible creasing to provide a product formed with a density between 40 and 400 kg/m³ preferably between 45 and 200 kg/m³ and more preferably between 50 and 150 kg/m³ possibly having a hinge resistant to repeated closure/ opening cycles.
- 39. Combinations of partly finished product according to any of the Claims 1-32 in multilayer structures to form products of various geometry such as rolls, blocks and foam sheet of significant resilience, corner pieces, protective containers for use in the electrical domestic appliance sectors, or for electronic

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products, in the food sector, for pharmaceuticals, for design and furniture, for mail order or envelopes for couriers.

40. Combinations of materials according to Claims 1 to 32, with other supports to provide multilayers mixed with wood, paper, cardboard, non-woven fabric, woven fabric of natural or synthetic fibres, aluminium or other metals for use in the packaging sector.

Products formed according to Cialm 38, used principally in the food packaging sector and in particular as trays for foods with a lifetime of the order of 30 days for packaging meet, delry products, vegetables, eggs, fruit, display containers for glass, plastic or metal packages of small dimensions, containers for frast food such as containers for hamburgers, potato chips and similar products; multi compartment containers for fast food and meals.

42. Products formed according to Claim 38, used for hot and cold liquids as cups for coffee and drinks, containers for soup of the type used in Aslatic countries and for other products having a high liquid content for fast food and meals.

- 49. Products formed according to Claim 38, used as containers for objects of small weight such as multicompartment trays for portable telephones and small electrical domestic appliances, in particular, 30 with mechanical properties such as to avoid phenomena of abrasion encountered with containers of pressed paper etc.
- 44. Products according to Claim 38, in the form of trays of for wrapping meat in supermarkets which provide for the use of absorbent or super absorbent materials for eliminating the presence of blood, fitted directly into the sheet, applied to the surface or in intermodiate layers between two shells welded together or under the film which renders the tray impermeable.
- 45. Products formed according to Claim 38, in the form of containers for oven and microwave use possibly surface treated to avoid excessive weakening of the container by the effect of the removal of the water.

Fig.1

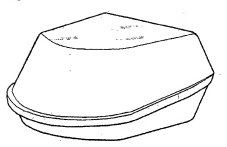
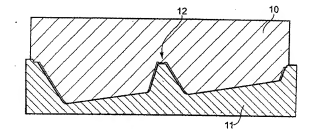


Fig.2







EUROPEAN PATENT APPLICATION

(88) Date of publication A3: 26.09.2001 Builetin 2001/39

(12)

(51) Int CI.7: **C08J 9/12**, B32B 5/18 // C08L3/00

(43) Date of publication A2: 29.08.2001 Bulletin 2001/35

(21) Application number: 01103390.9

(22) Date of filing: 14.02.2001

(84) Designated Contracting States: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL FT SE TR Designated Extension States: AL LT LY MK RO SI

(30) Priority: 15.02.2000 IT TO000141

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(54) Foamed starch sheet

(57) Materials in the form of foam sheet comprising destructured or complexed starch expanded as a continuous phase, having a density lying between 20 and 150 kg/m³, cell dimensions in the range lying between 25 and 700  $\mu m$  and with a cell distribution such that 80% of them have a dimension lying between 20 and 400  $\mu m$  in the absence of stretching.



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Application Number EP 01 10 3390

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